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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO
10/055,569	10/26/2001	Esha A. Gangolli	21402-191 (CURA 491)	8659
7590 09/22/2006		EXAMINER		
Jenell Lawson			PAK, MICHAEL D	
Intellectual property CuraGen corporation			ART UNIT	DA DED MINADED
555 Long Wharf Drive			ARTUNIT	PAPER NUMBER
New Haven, CT 06551			1646	
		DATE MAILED: 09/22/2006		

Please find below and/or attached an Office communication concerning this application or proceeding.

		Application No.	Applicant(s)	
Office Action Summary		10/055,569	GANGOLLI, ESHA A.	
		Examiner	Art Unit	
		Michael Pak	1646	
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A SHOR WHICHI - Extensio after SIX - If NO per - Failure tc Any reply	RTENED STATUTORY PERIOD FOR REPLY EVER IS LONGER, FROM THE MAILING DA ns of time may be available under the provisions of 37 CFR 1.13 (6) MONTHS from the mailing date of this communication. riod for reply is specified above, the maximum statutory period w or reply within the set or extended period for reply will, by statute, y received by the Office later than three months after the mailing atent term adjustment. See 37 CFR 1.704(b).	ATE OF THIS COMMUNICATION 36(a). In no event, however, may a reply be tirr vill apply and will expire SIX (6) MONTHS from cause the application to become ABANDONE	N. hely filed the mailing date of this co D (35 U.S.C. § 133).	
Status				
2a)⊠ Th 3)⊡ Si	esponsive to communication(s) filed on <u>23 Ju</u> nis action is <b>FINAL</b> . 2b) This note this application is in condition for allowards and in accordance with the practice under E	action is non-final. nce except for formal matters, pro		e merits is
Disposition	of Claims			
4a 5)⊠ CI 6)⊠ CI 7)□ CI	aim(s) 5-7,9,12-14,30,33 and 43-48 is/are per of the above claim(s) is/are withdraw aim(s) 5-7, 9, 12-13, 30, 33 is/are allowed. aim(s) 14 and 43-48 is/are rejected. aim(s) is/are objected to. aim(s) are subject to restriction and/or	vn from consideration.		
Application	Papers			
10)∐ The Ap Re	e specification is objected to by the Examiner of drawing(s) filed on is/are: a) acception and request that any objection to the objectement drawing sheet(s) including the correction of the coath or declaration is objected to by the Examiner.	epted or b) objected to by the Edrawing(s) be held in abeyance. See on is required if the drawing(s) is obj	e 37 CFR 1.85(a). ected to. See 37 Cl	
Priority und	ler 35 U.S.C. § 119			
12) Acl a) 1. 1. 2. 3.	knowledgment is made of a claim for foreign  All b) Some * c) None of:  Certified copies of the priority documents  Copies of the certified copies of the priority documents  population from the International Bureau  the attached detailed Office action for a list of	s have been received. s have been received in Application ity documents have been received (PCT Rule 17.2(a)).	on No ed in this National	Stage
Attachment(s)	f References Cited (PTO-892)	4) 🔲 Interview Summary	(PTO_412)	
2)  Notice of 3)  Informati	f Draftsperson's Patent Drawing Review (PTO-948) con Disclosure Statement(s) (PTO/SB/08) b(s)/Mail Date	Paper No(s)/Mail Da 5) Notice of Informal P 6) Other:	ite	

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#### **DETAILED ACTION**

### Response to Amendment

- 1. Amendment filed June 23, 2006 has been entered.
- 2. The text of those sections of Title 35, U.S. Code not included in this action can be found in a prior Office action.
- 3. Applicant's arguments filed June 23, 2006, have been fully considered but they are not found persuasive.
- 4. Claims 1-4, 8, 10-11, 15-29, 31-32, and 34-42 are cancelled. Claims 5-7, 9, 12-14, 30, 33 and 43-48 are pending.

#### Claim Objections

5. Claims 43-48 are objected to under 37 CFR 1.75(c), as being of improper dependent form for failing to further limit the subject matter of a previous claim.

Applicant is required to cancel the claim(s), or amend the claim(s) to place the claim(s) in proper dependent form, or rewrite the claim(s) in independent form. Claims 43-48 are broader than than the dependent parent claim 5. It is suggested that claims refer to SEQ ID NO:19 instead of the claim 5.

## Claim Rejections - 35 USC § 112

6. Claims 43-48 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention. This is a written description rejection.

The reason for the rejection has been set forth in the previous office action.

It is suggested a functional language be used to limit the variants to enable nucleic acids encoding the protein.

Newly amended claims 43-48 encompass an isolated nucleic acid variants. However, the essential feature of the invention is the nucleic acid encoding the specific polypeptide of SEQ ID NO:20. The sole utility and function of the nucleic acid is the diagnostic screen for cancer, and one of skilled in the art cannot envision the full genus of variant nucleic acid which can be used for diagnostics. The claims encompass variants whose structural change cannot be used to diagnose cancer cells versus normal cells since the change in the nucleic acid will cause the loss of the diagnostic capability of the probe. Claimed nucleic acid variants encompass a large genus which are alleles or variants whose function has yet to be identified from different species of animal because the structure of the newly identified naturally occurring protein is not known. *University of California v. Eli Lilly and Co. (CAFC) 43 USPQ2d 1398* held that a generic claim to human or mammalian when only the rat protein sequence was disclosed did not have written description in the specification.

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7. Claim 14 remains rejected under 35 U.S.C. 112, first paragraph, because the specification, while being enabling for isolated cells transfected or transformed with an expression vector, does not reasonably provide enablement for a cell comprising a vector. The specification does not enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the invention commensurate in scope with these claims.

The reason for the rejecton has been set forth in the previous office action.

It is suggested that the term "isolated" be inserted between the terms "cell" and "comprising".

The first paragraph of § 112 requires that the patent specification enable "those skilled in the art how to make and use the full scope of the claimed invention without 'undue experimentation." Genentech, Inc. v. Novo Nordisk AIS, 108 F.3d 1361, 1365, 42 USPQ2d 1001, 1004 (Fed. Cir. 1997) (quoting In re Wright, 999 F.2d 1557, 1561, 27 USPQ2d 1510, 1513 (Fed. Cir. 1993)); see also In re Fisher, 427 F.2d 833, 839, 166 USPQ 18, 24 (CCPA 1970). ("[T]he scope of the claims must bear a reasonable correlation to the scope of enablement provided by the specification to persons of ordinary skill in the art."). Whether making and using the invention would have required undue experimentation, and thus whether the disclosure is enabling is a legal conclusion based upon several underlying factual inquiries. See In re Wands, 858 F.2d 731, 735, 736-37, 8 USPQ2d 1400, 1402, 1404 (Fed. Cir. 1988). As set forth in Wands, the factors to be considered in determining whether a claimed invention is enabled

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throughout its scope without undue experimentation include the quantity of experimentation necessary, the amount of direction or guidance presented, the presence or absence of working examples, the nature of the invention, the state of the prior art, the relative skill of those in the art, the predictability or unpredictability of the art, and the breadth of the claims.

Likewise, in Amgen Inc. v. Chugai Pharm. Co., 927 F.2d 1200, 18

USPQ2d 1016 (Fed. Cir. 1991), the court affirmed the holding of invalidity of claims to analogs of the EPO gene under § 112 for lack of enablement where applicants had claimed every possible analog of the EPO gene but had disclosed only how to make EPO and a very few analogs. "[D]espite extensive statements in the specification concerning all analogs of the EPO gene that can be made, there is little enabling disclosure of the particular analogs and how to make them .... There may be many other genetic sequences that code for EPO-type products. Amgen has told how to make and use only a few of them and is therefore not entitled to claim all of them." Id., 927 F.2d at 1213-14, 18 USPQ2d at 1027.

Claims encompass cells comprising a vector which has not been isolated and includes cells from gene therapy or transgenic animal. However, one skilled in the art cannot make and use cells for gene therapy or transgenic animal because such techniques require undue experimentation. The state of the art is such that one skilled in the art must determine the phenotype of a transgenic animal. The amount of direction provided in the specification is limited to what is practiced by one skilled in the art, which is to transfect isolated cells using nucleic acid, subcloned into a vector. One

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skilled in the art would require empirical experimentation in order to determine how to create a transgenic animal or gene therapy. Such experimentation is unpredictable and requires undue experimentation. Therefore, based on the above <u>Wands</u> analysis, a preponderance of the evidence supports a conclusion that one skilled in the art would not have been enabled to make and use the claimed invention without undue experimentation.

- 8. Claims 5-7, 9, 12-13, 30 and 33 are allowed.
- 9. Applicant's amendment necessitated the new ground(s) of rejection presented in this Office action. Accordingly, **THIS ACTION IS MADE FINAL**. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the date of this final action.

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10. Any inquiry concerning this communication or earlier communications from the

examiner should be directed to Michael Pak whose telephone number is 571-272-0879.

The examiner can normally be reached on 8:30 - 2:00.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's

supervisor, Gary Nickol can be reached on 571-272-0835. The fax phone number for

the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the

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USPTO Customer Service Representative or access to the automated information

system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

Michael Pak

**Primary Patent Examiner** 

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17 September 2006